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#### Constitutively Active Receptors

				•									2H2C_rat	nemuhu Vztac	C113 A C113	H1B_human	CLASS A GROUP II	į.	MSHR_mouse	CLASS A GROUP I	File Name
	*												5-hydroxytryptamine <sub>2C</sub>	3-ilydroxytryptammc2A	5 hudovutantomine	5-hydroxytryptamine <sub>18</sub>		MSH	melanocyte-stimulating horimone	4	Receptor
													C-terminus of IC3	(	C-terminus of IC3	C-terminus of IC3			TMII		Mutation Site
												Ľ	312 NEDDA <u>S</u> KVLGI	NEQKA <u>C</u> KVLGI K	322	313 RERKAIKTLGI K,R,Q			VSIVLETTIIL  K		Sequence
					1.	•							PI hydrolysis / COS-7		IP production / COS-7	binding of ["S]GTP[S]/			HEK293, stably transfected		Assay / Cells
-			4					1					al. 1997)	al. 1998)	(Egan, Herrick-Davis et	(Patwels, Gouble (1999)			(KODOIDS, Nadeau et al. 1993)		Reference

:

ACM2-numan		ACMI_human	A2AA_human	\1AB_human	AIAB_human			AlAB_human	AIAB_human		8	CLASS A
IIIuscal IIIIc accy colored	muscarinic acetylcholine M1	muscarinic Hm1	α <sub>2</sub> C10-adrenergic	α <sub>1B</sub> -adrenergic	α <sub>1B</sub> -adrenergic			α <sub>18</sub> -adrenergic	α <sub>1B</sub> -adrenergic		α <sub>ιΒ</sub> -adrenergic alpha 1B-AR	
	junction of IC3 and TMVI	C-terminal IC3 loop junction	C-terminal IC3 loop	C-terminus IC3	C-terminal IC3	TMV	carboxyl end of IC3	TMIII	junction between TMDIII and IC2	junction between TMDIII and IC2	TMDI	
KKVTRTIL <sub>1</sub> A 1-4 A inserted	390	360 SLVK <u>E</u> KKAARTLS	373 (3487) EKRF <u>T</u> FVLAV X=F,A,C,E;K	288 293 KFSREKKAAKTLGI K H L	293 SREKKAAKT X=19 different substitutions	EEPFYALFSSLG V	293 REKKAAKTLGI E	128 AVDVL <u>C</u> CTASI F	143 CAISIDRYIGV K	142 CAISI <u>D</u> RYIGV A	63 FAIVGNILVIL A	-
COS-7	IP production, inhibition	P1 / HEK(UZ93)	inhibition / HEK 293	PI hydrolysis / rat fibroblast	PI / COS-7	IP / COS-1	IP arachidonic acid release	IP / COS-1	IP / COS-7		IP/COS-7	
	(Liu, Blin et al. 1996)	1995)	(ACM, ACCOUNTS Shockley et al.	(Allen, Leinowitz v 1991)	(Kjelsberg, Cotecchia et al. 1992)	(Hwa, Galvin et al. 1997)		(Perez, riwa et al. 1990)	(Scheer, Costa et al. 2000)		(Scheet, Fancin et al. 1997)	

ion / (Cho, Taylor et al. 1996)		N, A		\$	
on/	IIIIIN	N, A		-	
	cAMP production	115 FMISL <u>D</u> RYCAV	IC2	histamine H <sub>2</sub>	HH2R_rat
	COS-7	286 FVCCWLPFFIL A	IVMI	dopamine D1	DADR_human
_	CAND	288 from D1B receptor APDTSIKKETKVLKT			<del></del>
cAMP (Charpentier, Jarvie et al. EK293   1996)	adenylyl cyclase; cAMP accumulation / HEK293	264 SFKMS <u>F</u> KRETKVLKT I K	carboxyl terminal IC3	dopamine DIA	DADR_human
	activation; agonist binding affinity / COS-7 or CHO	X EEK	C-ICHIIIIIai ICJ ROOP	β <sub>2</sub> -adrenergic beta-2AR	B2AR_human
(Samama, Cotecchia et al.	adenylyl cyclase	266 272	C to lead to lead		
agonist (Mason, Moore et al. 1999)	adenylyl cyclase; agonist binding / CHW	389	C-terminus	β <sub>1</sub> -adrenergic	BIAR_human
	binding / NIH-3T3	YNIMVLVSTFCDKCV X=V,F,R,K,+more	junction of IMVI and ECS	m5 muscarinic muscarinic acetylcholine M5	ACM5_human
(Spalding, Burstein et al.	R-gal radioligand	T V		muscarinic acetylcholine Mo	
(Spalong, Durstem et al.	β-gal; radioligand binding / NIH-3T3	451 459 465 AILLA FIITW TPYNI MVLVST M L H C	IVMI	m5 muscarinic	ACM5_human
	<del>                                     </del>	45 466	TMVI	muscărinic acetylcholine M5	
(Burstein, Spaiding et al.	β-gal/NIH 3T3	chimera composed of m2 1-69	N-terminus to TMII	m5 muscarinic	ACM5_human
1994)	IP / COS-/	TWTPYNIMVLVNT S	IMVI	m3 muscarinic (rat). muscarinic acetylcholine M3	ACM3_rat
(Blin-I Water				- K	CLASS A GROUP II

LOOPART LOESEN

			. <u>)                                    </u>	OPSD_human		QPSD_human		OPSD_human				OPSD_human	CLASS'A GROUP III	ਨ -	
	47		rhodopsin	opsin		opsin	rhodopsin	opsin			шофран	opsin		Receptor	
	IC2			IIVMI	plus TM3	TM6		IIIMI		IWVII	TMIII	IIMI		Mutation Site	
	134 VVLAI <u>E</u> RYVVV Q	disrupts critical salt bridge between <sup>296</sup> Lys(TMVII) and <sup>113</sup> Glu(TMIII)	X=E,M natural mutants + 10 different a.a. substitutions	PAFFAKSAAIY	plus G113Q	257 RMVIIMVIAFL Y,N	I,Q,S	134 VVLAI <u>E</u> RYVVV	hat cut	292 296 MTIPAFFAKSAAIY E G,E,M	113 GCNLEGFFAT Q	90 PMVLGGFTSTLY		Sequence	
		~		transducin; radioligand binding / COS		transducin, GTP/S uptake / CQS		transducin; radioligand binding / COS				transducin; phosphorylation by rhodopsin kinase / COS		Assay / Cells	
	(Cohen, Yang et al. 1993)		(Concil, Lang Stat. 1999)	(Govardnan and Oprian 1994);		(Han, Smith et al. 1998)		(Acharya and Karnik 1996)				(Kum and Oprian 1993)		Reference	

100	Marie Company			•		TRFR_mouse		
, S.	· Brown			•	TRH-R	in-releasing hormone		
	A CONTRACTOR OF THE PROPERTY O		1000年の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の日本の		11 11 11 11 11 11 11 11 11 11 11 11 11	carboxyl tail		
	1、1、1、1、1、1、1、1、1、1、1、1、1、1、1、1、1、1、1、	語が、中心の変数を含っている語数の音響であった。 ・アインをできる音楽をおける音響をよっている。	が、これできる時間ではなった。これは、では他のではないとしています。 では、これでは、ないのでは、これでは、これでは、これでは、これでは、これでは、これでは、これでは、これ		dOLS	FRKICNCKOK		
				stably transfected	IP formation / AtT20,	Xenopus oocytes;	1 (50° 3+ *WILL (Ca2+)	
	一般は、 は、 は、 はないないない。					<u>=</u>	(Matus-Leibovitch,	第二年の1日本の1日本の1日本の1日本の1日本の1日本の1日本の1日本の1日本の1日本

				BRB2_human	GROUP IV	CLASSA		-	
	•	BK-2		bradykinin B <sub>2</sub>			Receptor		×
			IMVI	ТЩШ		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Mutation Site		
	শ্য	LLFIICHLPFQI	A	AIISMALYSSI	113		Schence	2000000	
			•		IP production / COS-7 (Marie, Koch et al. 1999)			Assay / Cells	
				•	(Marie, Koch et al. 1999)		**************************************	Reference	

	OXYR_human		OPRD_mouse		LSHR_rat		LSHR_human	·	LSHR_human		LSHK_numan	T CUB human		IL8B_human				FMLR human		AG2R_rat				GROUP V	File Name	4.
	oxytocin		delta opiod receptor	(LH/hCG)	luteinizing hormone / human		luteinizing hormone (LH)		luteinizing hormone (LH)		Inchirzing morning ()	Inteinizing hormone (LH)	CXCR-2 chemokine	interleukin-8 receptor B			anine (fMLPR)	formylmethionylleucylphenylal	Type-1 A angiotensis II	ÄΤ <sub>ικ</sub>		Typę, 1A angiotensis II	ATIA		Receptor	外意义
7		103	TM3		TMVI		TM6		TA AT	TWVI		IC3		IC2				ICI	other multiple mutations	C-terminus of 1M/					Ivation City	Mutation Site
	LMSLDRCLAIC A	137	KVLS I X K. H	G, Y	ILIFTDFTCMA		KIAKKMAILIFIDETCM		ILIFTDFTCMA	578	MATNKDTKIAKK	564	•	ACISVDRYLAIVH	SWISS-PROI Cacabase/	(K above conflicts with	LYVWVTAFEAKRTINAIWFLNLAVA	51	Q	TEAGETOKKEK	TAL (TMATT) THE CETTOR	6	ASVSTALYASV			Sequence
		IP production / COS-7	inhibition / COS-7	adenylyl cyclase	HEK 293T	ca MP production /	cos-7	caMP production /	COS-7	cAMP production /	HEK293	cAMP production /	NIH 3T3	moblization and actin	IP production; Ca2+		stimulation / COS-7	phospholipase C	mobilization / CIA	293; intrcellular Ca2	IP production / HEK-	の対する。 では、「大きなのでは、大きなのでは、大きなのでは、「大きなのでは、大きなのでは、大きなのでは、大きなのでは、「大きなのでは、「大きなのでは、「大きなのでは、「大きなのでは、「大きなのでは、	IP production / CQS-7.	phospholipase C;		Assay / Cells
	1999)	(Fanelli, Barbier et al.	1999)	(Cavalli, Babey et al.	1997; Bradbury and	(Bradbury, Kawate et al.	1995)	(Kosugi, Van Dop et al.	1993)	(Shenker, Laue et al.		(Kudo, Usuga et al. 1770)	1 1000	1999)	(Burger, Burger et al.			Graonic et al. 1995)	(Amaruda, Draga,	2000)	(Parnot, Bardin et al.		al. 1997)	(Groblewski, Maigret et	を できる	Reference

Contract Balance Balance

#### TOSZOT SHOKEDS

	TSHR_human		TSHR_human	•	TSHR_human		•		TSHR_human			•		TSHR_human		TUDD human			RE23_mouse		*	PE23_human		PAFR_human		PAFR_human	
thyroid stimulating hormone	thyrotropin (TSHR)	thyroid stimulating hormone	thyrotropin (TSHR)	thyroid stimulating hormone	thyrotropin (TSHR)			thyroid stimulating hormone	thyrotropin (TSHR)				thyroid stimulating hormone	thyrotropin (TSHR)	THE CHICAL	thrombin		EPS	prostaglandin E <sub>3</sub>		EP3III	prostaglandin E3,		platelet-activating ractor (rAr)	(DAE)	platelet-activating factor (PAF)	-
	IC3		IIVMT		TMV		TMVII		TMIII		EC2	# C		BC1	•	EC2 loop	-		carboxyl-terminal tail			C-terminal tail		TIME		C-terminus of IC3	
gerecton	VRNPQ <u>YNPGDKDTK</u> IAK		677 CANPFLYAIFT	L	VAFVIVCCCHV	KI	YPLNSCANPFL	;	ASELS <u>V</u> YTLTV		YAKVSICLPMD	568	F, M	486 YYNHAIDWQTG	CHDVLNETLLEGYYAYY DLKO KDF I	259 268	cat	MMNHL (3B)	.336 (3α)	îtruncated	FCOMRKERLEGOESFWGN	360 BCORREWGN	A	CLFFINTYCSV	100	EVKRRALWMVCTVLAV	3 3 3 4
·	COS-7	AND formation /	CHO cells	AND famation /	cAMP formation '. <sub>1</sub> COS-7 cells				activation / COS-7	adenuly cyclase			COS-7	inositol phosphate diacylglycerol cascade /	hydrolysis, reporter gene induction / COS-7	45Ca 2+ efflux, PI		expressed	cyclase / CHO, stably			cyclase / CHO-K1	cylcase inhibition / CHO	production, adenylyl	arachnidonate release, IP	r production of the	IP production / COS-7
	al. 1998)	(Wonerow Schoneberg et	The state of the s	(Russo Wonder al. 1999)	(ESapa, Dupiez et al. 1999)	3			1994)	(Duprez. Parma et al.				(Parma, van Sange et al. 1995)	1996)	(Nanevicz, Wang et al.			1996)	Minney Nerichi et al		(JIII, Mao et at. 1777)	1007		(Ishii, Izumi et al. 1997)	1996)	(Parent Le G uill et al.

DGZOT SHAKEOUT

V2R_human vasopressin V2	thyroid sti	TSHR_human thyrotropin (TSHR)
	thyroid stimulating hormone	
ß		IC3/TMVI
136 LAMTL <u>D</u> RHRAI A	V	623 632 KDTKIAKRMAVLIFIDFICM
COS-7		cAMP activation / COS-7
	(Morin Cotte et al. 1998)	(Paschke, Tonacchera et al. 1994)

Figure 1 (Page 9 of 15)

#### TOSECT SHESTOOT

	VIPR_human		ĠLR_rat		CLASS B			B	CLASS B GROUP II	CALR_human h	_	File Name   R	
**	vasoactive intestinal peptide 1 (VIP)		glucagon	glucose-dependent insulinotropic peptide (GIP-R)				parathyroid hormone PTH / PTH-related peptide	<i>b</i> /.	human calcitonin hCTR-1 hCTR-2		Receptor	
junction of IC loop 3 and TMVI	junction of IC loop 1 and TMII	IC end of TMVI	junction of IC loop1 and TMII	IVMI			junction of IC3 and TMVI	junction of IC1 and TMII		wild type (native) protein		Mutation Site	
343 LARSTLLLIP X= K, P	178 RNYIHMHLFI requires functional integrity of the N-terminal EC domain	352 RLARSTLTLIP A	178 TRNYI <u>H</u> GNLFA R	340 VFAPVIEEQAR P			410 KLLKSTLVLMP C,others	TRNYIHMHLFL R, K				Sequence	
	cAMP production / COS-7 or CHO		COS-7	cAMP production / L293	1700			COS-7	Total I	adenylyl cyclase cAMP production / COS-1		Assay / Cells	
¥	(Gaudin, Maoret et 1998) (Gaudin, Rouyer-Fessard et al. 1998)		(1998)	(1seng and Lin 1997)	(Tone and I in 1997)			1997)	(Cohinani Jensen et al	(Concn, Inaw et al. 1997)	1 1007)	Kelerence	J-1

					1	Cartan State of State	•	CASR_human	CLASS C	Life Maine	Tile Nome		
								calcium-sensing	4 3	Merephy.	Decement		**************************************
								N-terminal EC		THE PARTY OF THE P	Mutation Site		
							multiple combinations	various substitutions, in	TI.SPVAONKIDSLNLDEPCNCSBHI		Sedhence	800000000000000000000000000000000000000	
•								•	IP/tsA			Assav / Cells	
		  - 						2000)	(Jensen, Spalding et al.		A AMERICAN IN THE PARTY OF THE	Reference	

Figure 1 (Page 11 of 15)

					5
File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Kelerence
CLASS D				The state of the s	Mischiele Rrown et al.
074283 RCB2	pheromone	TM6	PLSAYQIYLGT	neterologous yeast assay	1999)
C. cincreus			7	lac7 reporter gene	(Kononka Margarit et al.
STE2_yeast	pheromone α-factor	TM6	QSLLV <u>PS</u> IIFI	such schooms Bonn	1996)
		double mutations TMS	223	lacZ reporter gene /	(Dube, DeCostanzo et al.
SIEZ_yeast	pheromone a-tactor	COCCA International action	MSFVLYVKMILAIR	yeast	2000)
		and	C C		
		TM6	DSFHILLIMCQSLL		
			cc cc		
			double mutations double mutations		
STE3_yeast	pheromone a-factor	IC3	194 DVRDILHCINS	β-galactosidase	(Boone, Davis et al. 1993)
CTR2 veget	nheromone o-factor	TM6	253 258	β-galactosidase	(Sommers, Martin et al.
\$1 <i>52_</i> yeast	ристопноне се-таског	1	LIMSCQSLLV <u>PS</u> IIFI		2000)

Acharya, S. and S. S. Karnik (1996). "Modulation of GDP release from transducin by the conserved Glu134-Arg135 sequence in rhodopsin." L Biol Chem 271(41): 25406-11. Alewijnse, A. E., H. Timmerman, et al. (2000). "The Effect of Mutations in the DRY Motif on the Constitutive Activity and Structural Instability of the Histamine H(2) Receptor."

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#### A Point Mutation Enhances MC-4 Receptor cAMP production (% a-MSH stimulated) Constitutive Activity © α-MSH 1 μM

basa

Figure 2

MC-4 WT

D146M

MC-4

Light units Light Emission Induced by the WT CCK-BR pcDNA1 CCK-BR CCK-BR vs. a Constitutively Active Mutant SMS-Luc L325E pcDNA1 CCK-BR SRE-Luc CCK-BR L325E 四 başuı

Figure 3

# A Point Mutation Confers Constitutive Activity to the Rat $\mu$ Opiod Receptor

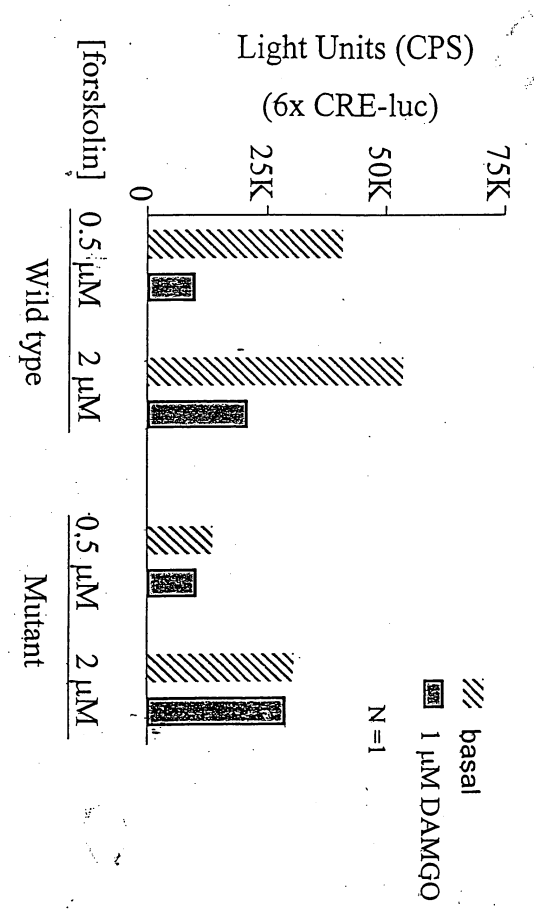
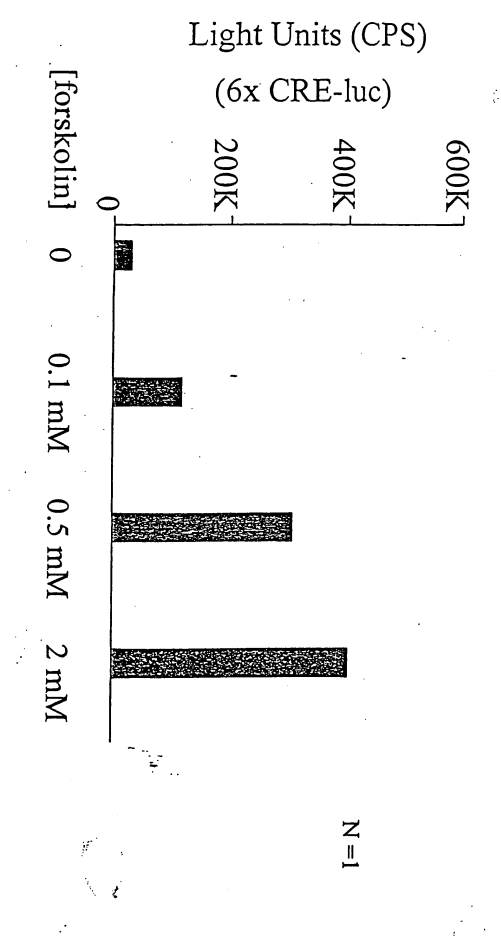


Figure 4

# Forskolin Stimulated HEK293 Cells Transfected With pcDNA1 and a CRE-luc Construct



# The Rat $\mu$ Opioid Receptor Signals Through Gai

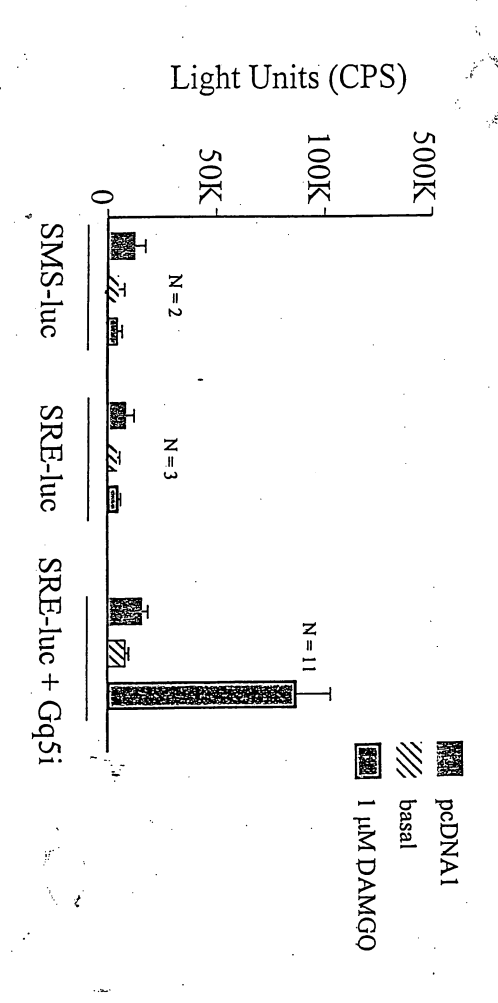


Figure 6

# Point Mutation Confers Constitutive Activity to the Rat $\mu$ Opioid Receptor

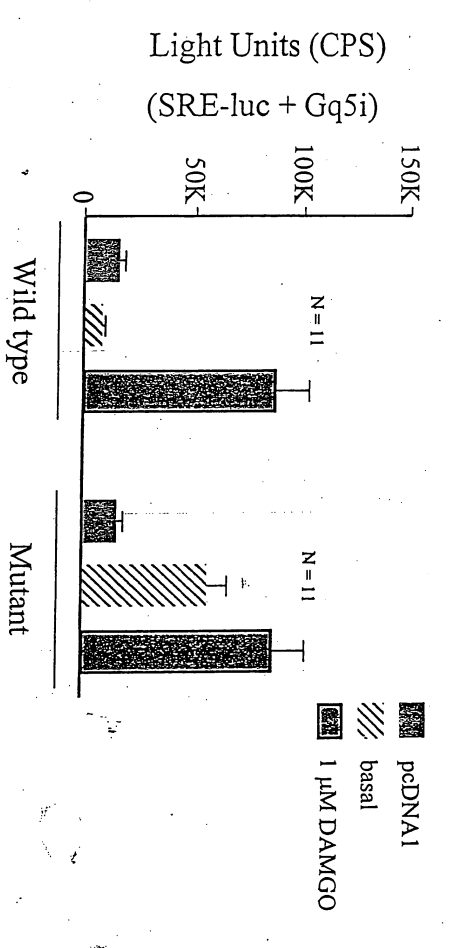
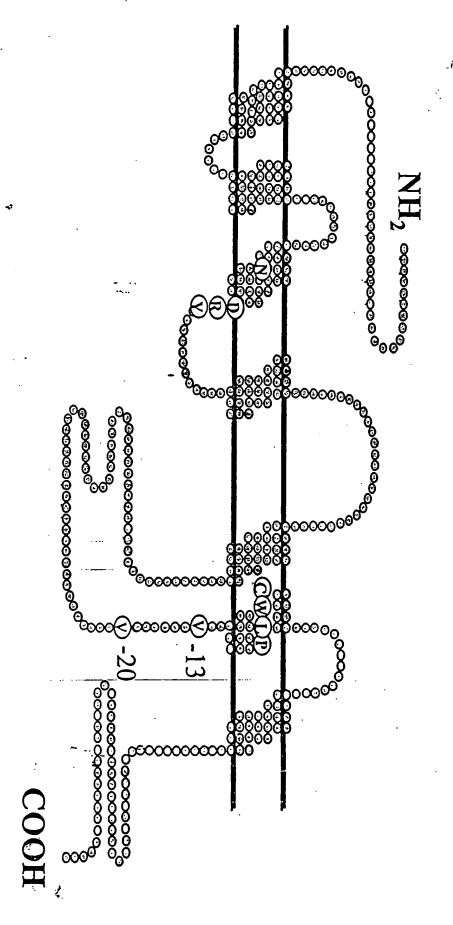
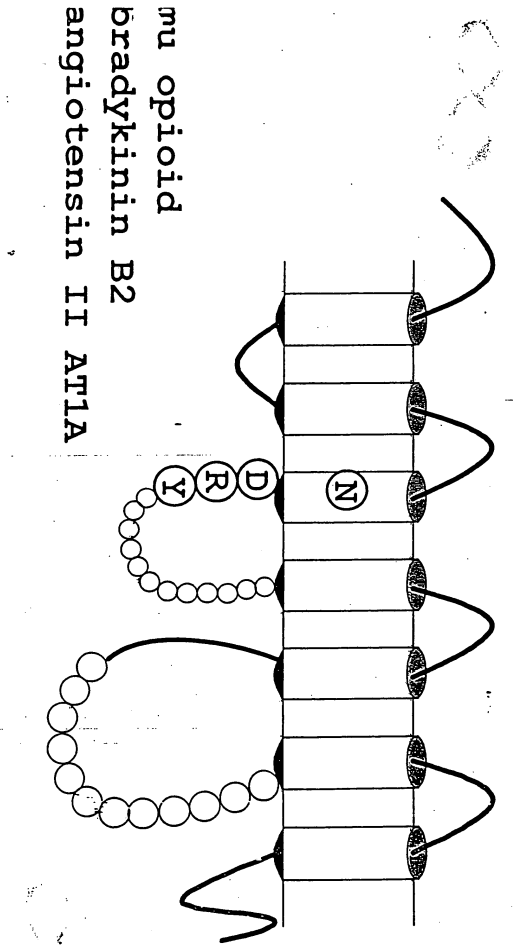


Figure 7

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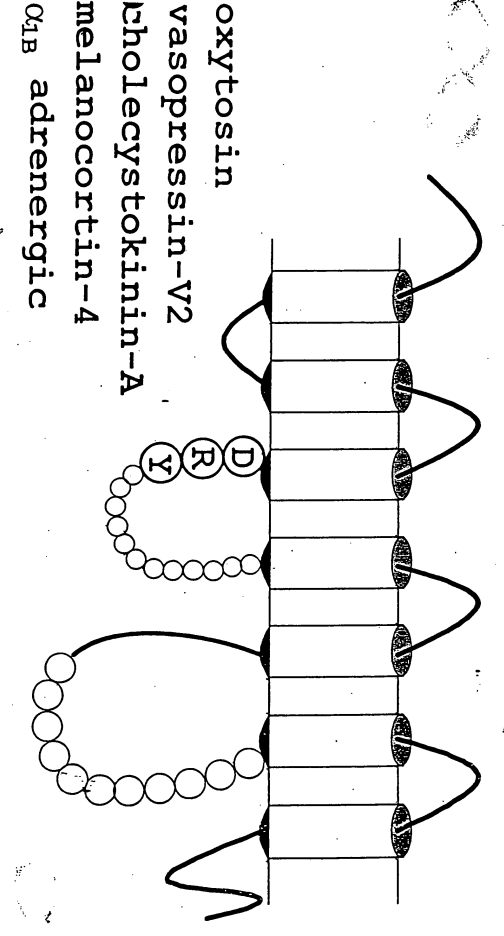


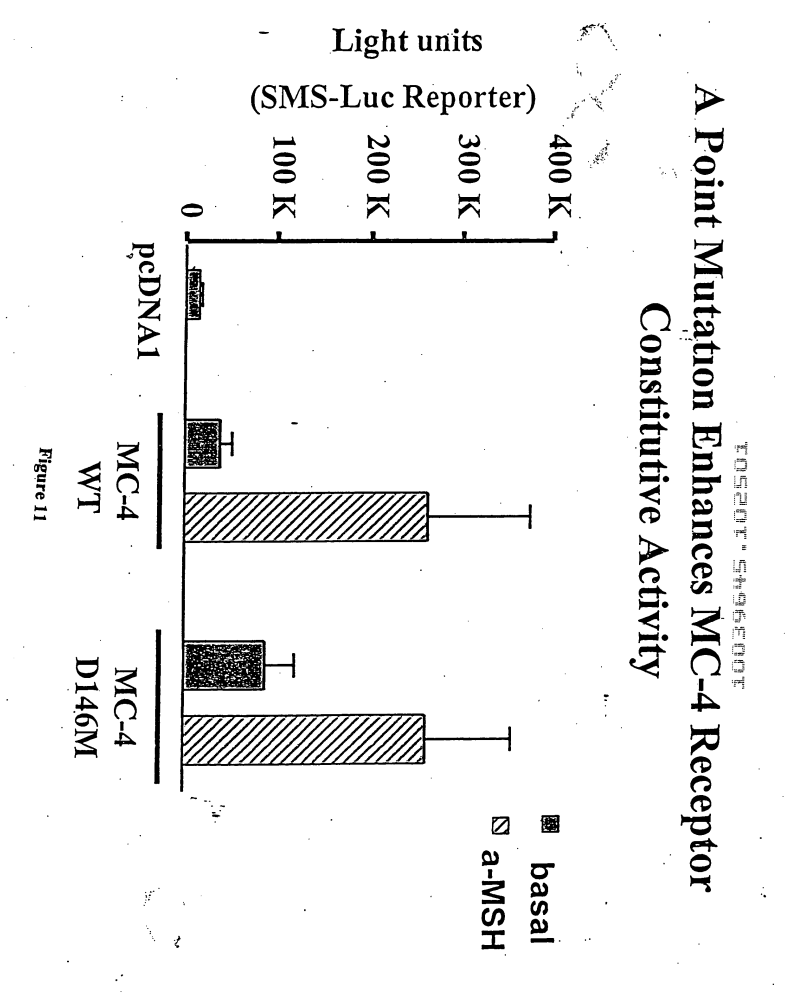
# for Mutation Induced Constitutive Activity TMD III Asn (-14 from DRY) is a Target



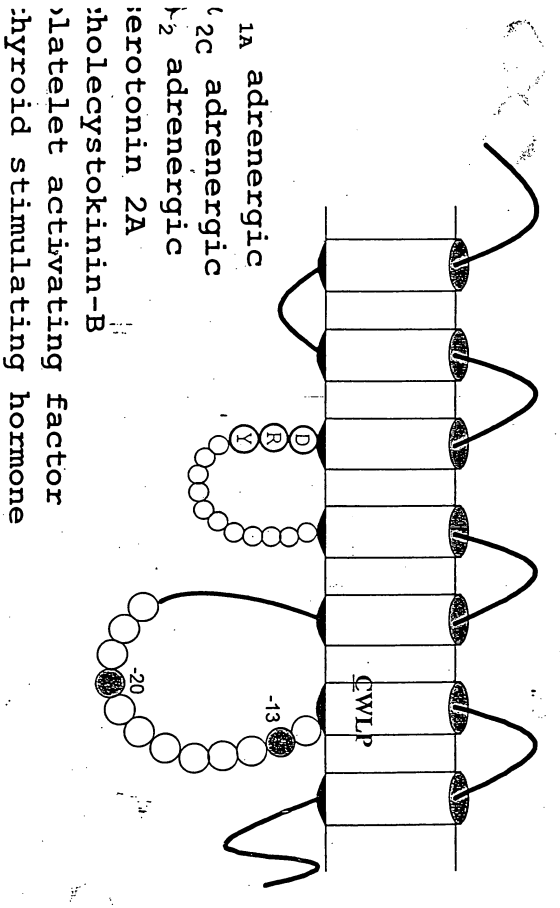
TOSZOT SHASEOT

# The 'DRY' Motif is a Target for Mutation **Induced Constitutive Activity**





## The -13 Position is a Target for Mutation Induced Constitutive Activity



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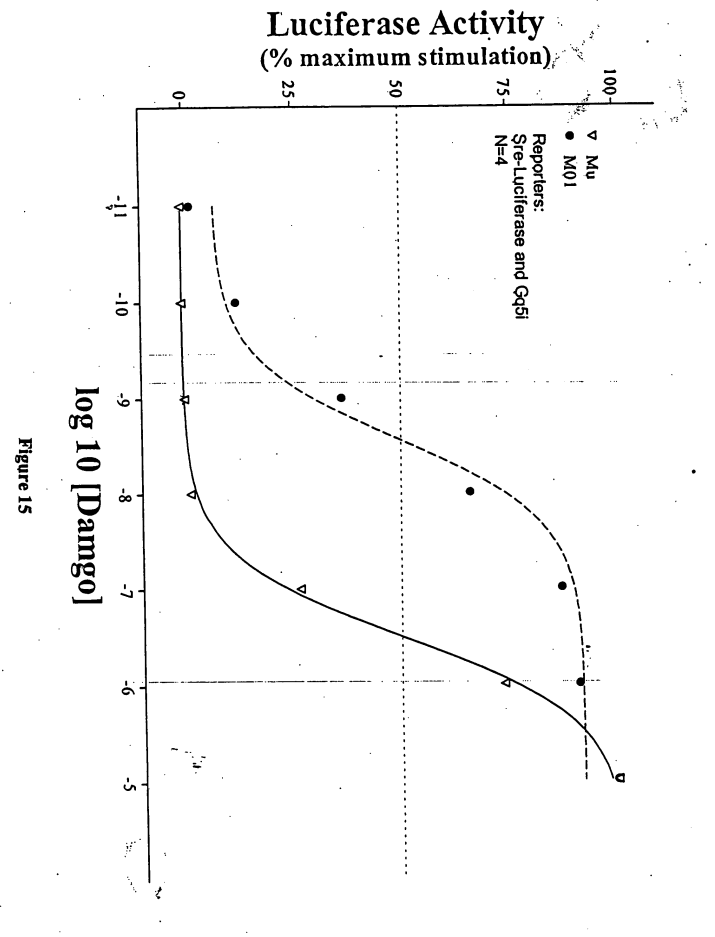
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## An Intracellular Point Mutation Results in Loss of Ligand-Induced Function

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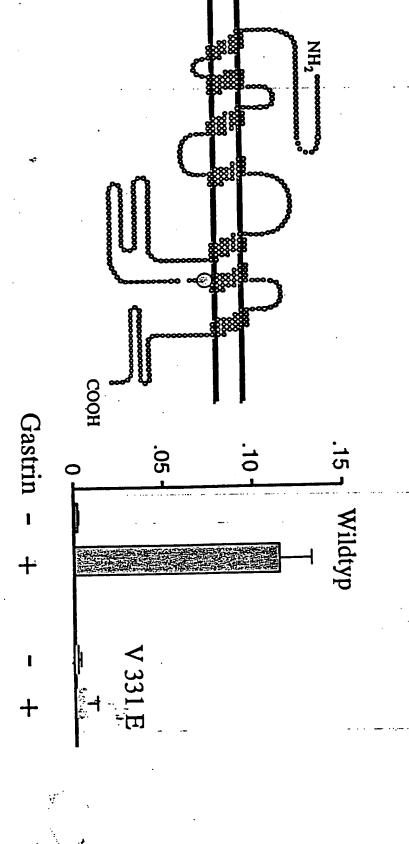


Figure 16

